LISTING OF THE CLAIMS

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1-47. (canceled)

- 48. (Previously presented) The method of claim 55, wherein said purified protamine fragment has a molecular weight of between about 400 and about 2000 Daltons.
- 49. (Previously presented) The method of claim 48, wherein said purified protamine fragment has a molecular weight of between about 500 and about 1350 Daltons.
- 50. (Previously presented) The method of claim 48, wherein said purified protamine fragment has a molecular weight of between about 1100 and about 1300 Daltons.
 - 51-54. (canceled)
- 55. (Previously presented) A method of inactivating heparin or low molecular weight heparin, comprising contacting heparin or low molecular weight heparin with a composition comprising an amount of at least a purified protamine fragment effective to inactivate heparin or low molecular weight heparin; wherein said purified protamine fragment is bioactive, has a molecular weight of between about 400 and about 2500 Daltons as determined by gel filtration and has reduced immunoresponsiveness or toxicity compared to native protamine.
- 56. (Previously presented) The method of claim 55, wherein said heparin or low molecular weight heparin is located within a mammal and said composition is administered to said mammal.
- 57. (Withdrawn) A method of ameliorating an effect of heparin or low molecular weight heparin in a mammal, comprising administering to said mammal at least a first pharmaceutical composition comprising an amount of at least a first purified protamine effective to ameliorate an effect of heparin or low molecular weight heparin in said mammal; wherein said purified protamine is bioactive, has a molecular weight of between about 400 and about 2500 Daltons and has reduced immunoresponsiveness or toxicity compared to native protamine.
- 58. (Withdrawn) A method for treating or preventing undue or excessive bleeding in a mammal, comprising administering to a mammal having or at risk for developing excessive bleeding at least a first pharmaceutical composition comprising an amount of at least a first

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purified protamine effective to treat or prevent undue or excessive bleeding in said mammal; wherein said purified protamine is bioactive, has a molecular weight of between about 400 and about 2500 Daltons and has reduced immunoresponsiveness or toxicity compared to native protamine.

- 59. (Previously presented) The method of claim 64, wherein said mammal exhibits excessive bleeding associated with systemic heparinization.
- 60. (Previously presented) The method of claim 64, wherein said mammal exhibits excessive bleeding associated with extracorporeal blood circulation.
- 61. (Previously presented) The method of claim 64, wherein said mammal exhibits excessive bleeding associated with a disease or disorder.
- 62. (Previously presented) The method of claim 64, wherein said mammal exhibits excessive bleeding associated with a trauma or surgery.
- 63. (Currently amended) The method of claim 64, wherein at least a coagulant is further administered to said mammal further comprising administering a coagulant to said mammal..
- 64. (Previously presented) The method of claim 56, wherein said mammal has or is at risk for developing excessive bleeding.
- 65. (Previously presented) The method claim 48, wherein said purified protamine fragment has a molecular weight of about 1300 Daltons.
- 66. (Previously presented) The method of claim 48, wherein said purified protamine fragment has a molecular weight of about 1200 Daltons.
- 67. (Previously presented) The method of claim 55, wherein said composition comprises at least a first and at least a second purified protamine fragment.
- 68. (Previously presented) The method of claim 56, wherein said mammal is a human subject.
 - 69. (Cancelled)

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70. (Previously presented) The method of claim 55 wherein inactivating heparin or low molecular weight heparin treats or prevents undue or excessive bleeding in a mammal.

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- [[70]] 71. (Currently amended) The method of claim 55 wherein the protamine fragment is a protease cleavage product.
- [[74]] 72. (Currently amended) The method of claim 70 wherein the protamine fragment is a protease cleavage product and said protease is selected from the group consisting of thermolysin, ficin, collagenase, kallikrein and proline-specific endoprotease.
- [[72]] 73. (Currently amended) The method of claim 55 wherein the protamine fragment is derived from a protamine selected from the group consisting of salmon protamine and clupine protamine.
- [[73]] 74. (Currently amended) The method of claim 55 wherein the protamine fragment comprises five or six arginine amino acid residues and 1 proline amino acid residue.
- [[74]] <u>75</u>. (Currently amended) The method of claim 55 wherein the protamine fragment comprises a minimum of six arginine amino acid residues.